

A METHOD OF RECYCLING AN EXHAUSTED SELENIUM FILTER MASS

This invention relates to a process for reclaiming spent selenium filter mass containing an inert material, following take-up of mercury by a substance which contains selenium and is present in the filter mass. Selenium is present in spent selenium filters, both in unspent active filter mass, mainly as elemental selenium, and in reacted filter mass as mercury selenides.

Selenium filters are being used for taking up gaseous elemental mercury, normally in small amounts, from gases and contain a filter mass consisting of an active substance of elemental selenium, selenium sulphide or some other active selenium compound which is capable of reacting with elemental mercury. The main portion of the filter mass consists of a carrier of an inert material, which may consist of silica, aluminium oxide, or a ceramic material. Such a filter, which has been in general use for various purposes for about thirty years, is disclosed in US 3 786 619 which also teaches a suitable method for making the filter.

In use of the filter, the active substance is gradually consumed, forming mercury selenide, and after operation over a period of time the performance of the filter will be reduced so much that its ability to take up mercury and perform the cleaning is no longer adequate. The filter mass is then replaced with fresh filter mass and the spent filter mass must be deposited in a suitable manner. Such deposition is costly in itself and also requires payment of waste tipping fees. In addition, the deposition often causes problems for several other reasons, i.e., environmentally, and both the carrier material and the active substance are of considerable value. As indicated initially, the used filter mass contains some unspent selenium-containing active substance, while the remainder of the selenium content of the active substance has reacted with mercury to form mercury selenide, which is a very stable compound.

Typically, the spent filter mass may contain 1 to 5 % by weight of Hg and may still contain 3 to 5 % by weight of unspent elemental selenium.

This being so, a reprocessing method for reclaiming the active substance and/or the carrier material is highly desirable. It is also desirable to be able to separate mercury contained in the filter mass from the inert carrier so that the amount of material that has to be deposited is as small as possible, e.g. to reduce the costs for the deposition.

Previously, methods have been proposed for processing mercury-containing waste at relatively high temperatures and in the presence of selenium to remove the mercury of the waste in the form of gaseous selenides. In such a method, primarily developed for used button-type batteries, which is described in EP 0655794, the batteries are treated in a rotary furnace at about 800°C in the presence of selenium to evaporate mercury in the form of selenide, thereby making the batteries harmless.

Such prior art methods for the destruction of mercury-containing materials are not useful or even feasible for processing spent selenium filter masses, particularly so if the filter masses are to be reclaimed, because capture of mercury existing as elemental mercury, Hg^0 , and as selenide, HgSe , will be problematic. Separation of the selenium for reclaiming will also be troublesome.

The object of the invention is to provide a process by which spent selenium filter masses can be reprocessed, in a manner that is both economically and environmentally acceptable, for reclaiming both the selenium and the filter mass. With such a process, a spent selenium filter mass can be cleaned and reused, while captured mercury contained in the filter mass can be separated and deposited in a suitable stable form.

To that end, the filter mass is treated in the steps which are set forth in the accompanying claims. In the course of the reclaiming process, the used and

spent selenium filter mass is first treated with a hydrogen peroxide solution, suitably having a concentration of about 50 %, for leaching out essentially all of the unspent active substance in the filter mass to form selenious acid. The resulting selenious acid is then separated and isolated. Then, this selenious acid is advantageously transferred to an installation for producing new selenium filter mass.

The filter mass that has been freed from the solution is then treated with aqua regia, preferably at an elevated temperature, for dissolving essentially all of the mercury selenide contained in the mass. The remaining filter mass, which is mainly formed of the inert carrier, remains essentially undissolved. The aqua regia solution with the content of mercury dissolved in it and selenium from the dissolved mercury selenide are separated from the filter mass and isolated.

Suitably, the aqua regia solution is heated and aerated to evaporate any excess of aqua regia. If it is desired to reclaim selenium from the solution, SO_2 is introduced into the solution after a partial neutralisation by suitable pH adjustment, resulting in precipitation of selenium as elemental selenium, Se(s) , that can be used to produce selenious acid for use, if desired, in the production of new filters. After further partial neutralisation of the solution, the mercury can be precipitated in the form of some poorly soluble compound other than selenide, such as sulphide. Thus, the solution can be reprocessed for reclaiming the selenium it contains, and the mercury content can also be obtained in a form that is suitable for deposition in an environmentally safe manner. Such reprocessing is not always possible or desirable, however, and, after neutralisation, the mercury and the selenium contained in the solution will then be precipitated as HgSe , so that mercury is taken care of for deposition in that very stable form. As indicated, it is preferred, however, to obtain the selenium for reuse and depose the mercury in a different stable form, such as sulphide.

Being now essentially free from the aqua regia solution and only containing inert material, the filter mass is washed and dried and then, like the previously separated selenious acid, forwarded to production of new selenium filter mass.

The invention will now be described as a preferred embodiment illustrated in the figure which is a diagrammatic flow chart of a process according to the invention.

The selenium filter mass is first fed to a tank for "Leaching 1", in which the filter mass is leached with about 50 % hydrogen peroxide, H_2O_2 , for leaching out selenium, that is, the unspent content of selenium, not bound to mercury, of the active filter mass. This leaching, which can be carried out in several steps, results in formation of selenious acid, H_2SeO_3 , which can be used as a starting material in the production of a selenium filter mass as is disclosed in the previously mentioned US 3 786 619. A solution containing selenious acid is thus obtained and forwarded to a purifying and filtering unit from which pure selenious acid is further transferred to an installation for the production of selenium filter masses.

The leach residue from "Leaching 1", that is, the remaining filter mass with its content of mercury selenide, is then subjected to another leaching step in "Leaching 2". In this leaching, which is carried out with aqua regia, that is, concentrated nitric acid mixed with three times as much of concentrated hydrochloric acid, the solid mercury selenide will be dissolved while forming a solution containing selenium and mercury in ionic form. Optionally, the leach tank may be heated so that the leaching takes place at an elevated temperature. The leaching is carried on until all or at least almost all of the mercury selenide content has been dissolved.

The solution with its dissolved content of mercury and selenium is separated from the leach residue, which is essentially formed by the inert carrier of the mass. The leach residue is withdrawn and washed with water and dried so that

a suitable carrier raw material for the production of filter masses is obtained, whereupon it is forwarded to an installation for such production.

The solution of aqua regia from "Leaching 2" is heated and blown with air for removing excess aqua regia, whereupon the solution is partially neutralised in two steps so that the selenium can first be precipitated with SO_2 while forming Se(s) , whereupon the precipitate of elemental selenium is separated from the solution. Using sodium sulphide or a different sulphide, selenium is then precipitated as HgS which can be separated and withdrawn for deposition.

After suitable purification, the precipitate of elemental selenium, Se(s) , from the neutralisation step can be brought to the installation for filter production.

The inventive step of the invention will now also be elucidated from an environmental and economical point of view.

It can be estimated that spent filter mass contains at most 125 kg of mercury per cubic metre. When transformed to HgS of a density of 8100 kg/m^3 , a volume of about 18 litres of fine-grained sludge is obtained. Even if small amounts of other substances should accompany the precipitate, the volume will not exceed 50 litres. That is, the volume of mercury-containing waste to be deposited will only be about 5 % of the original volume. Thus, the need for deposition is reduced to one twentieth when spent selenium filters are reprocessed in accordance with the invention. If the selenium of the mercury selenides formed is reclaimed, there are basically no selenium wastes and attendant losses.

The cost reduction obtained by the process according to the invention is substantial. A direct conversion based on the volume would thus result in a reduction of the deposition cost by as much as 95 %. The reuse of the cleaned carrier mass also means a great cost reduction.

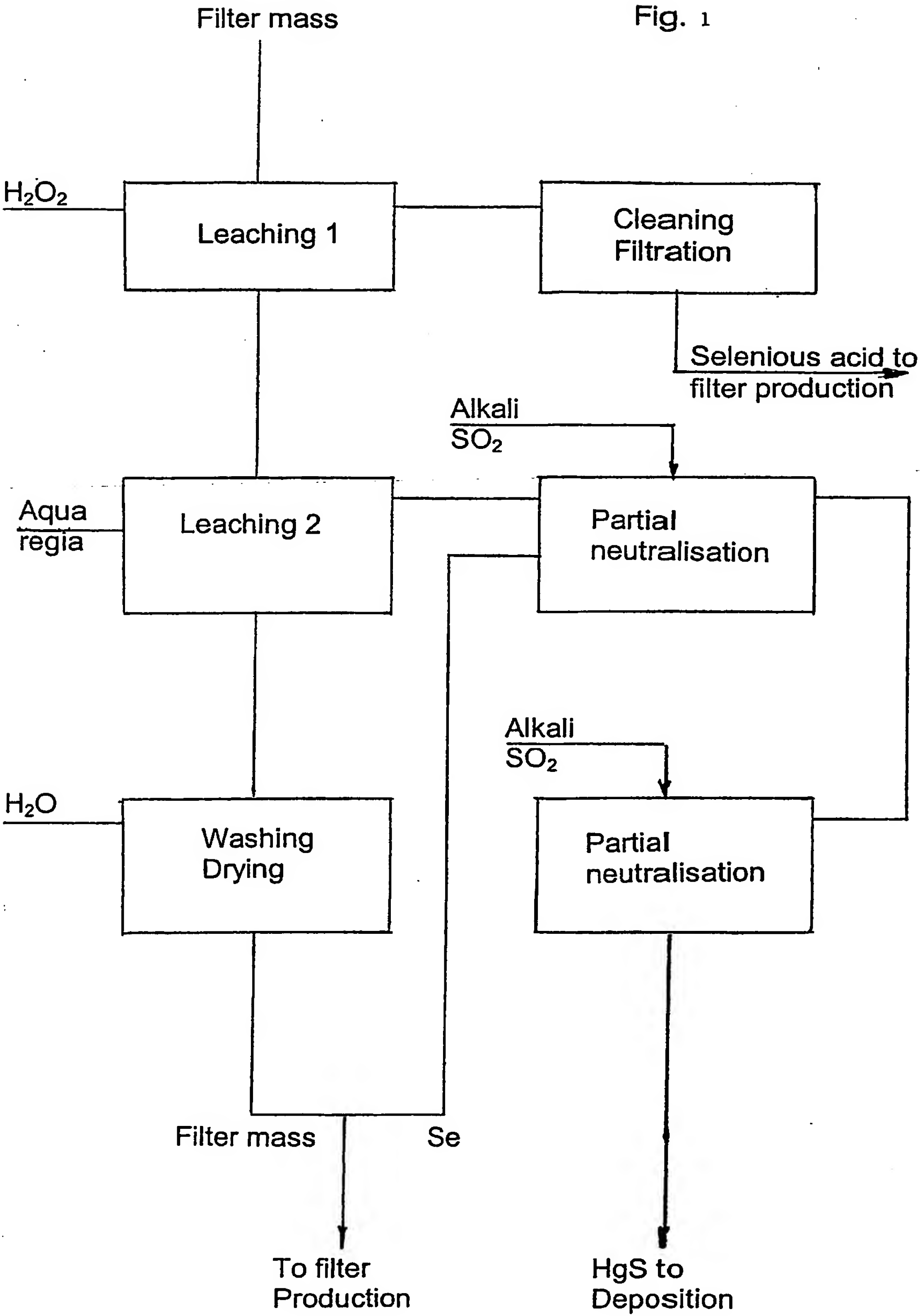
Example

In preliminary tests, leaching of the filter mass was carried out with an initial Hg content of 2.72 % and an initial Se content of 5.5 %. Leaching 1 using hydrogen peroxide resulted in leaching out of a total of 72 % of the selenium contained in the filter mass. In a following Leaching 2 using aqua regia, 99 % of the mercury content and also 25 % of the original selenium content could be leached out. The solution was neutralised, and selenium and mercury were precipitated as sulphides. After the precipitation the solution contained <0.001 % of the original content of mercury and 2.5 % of the original content of selenium. After washing of the leached mass, the mass contained only 0.2 % Hg of the original content, corresponding to a mercury content in the fully leached mass of about 0.005 %. These tests thus show that it is possible selectively to leach out the selenium which can then be returned to production of selenium filter masses. The leaching out of mercury has also been found to be effective, and the process according to the invention has reduced the volume to be deposited as mercury sulphide to about one eightieth of the original volume.

Claims

1. A process for reclaiming spent selenium filter mass containing an inert material, following take-up of mercury by a substance which contains selenium and is present in the filter mass, **characterised** in that the mass is treated with a hydrogen peroxide solution for leaching out of the selenium content in essentially all unspent active substance present in the mass to form selenious acid, in that the resulting selenious acid is separated and isolated for use, in that the mass freed of the solution is treated with aqua regia for dissolving essentially all of the mercury selenide present in it, in that the aqua regia solution with its content of mercury and selenium is separated from the mass and isolated and in that the mercury contained in the solution is precipitated and isolated in depositable form.
2. A process according to claim 1, **characterised** in that selenium is selectively separated and obtained from the aqua regia solution after adjustment of its pH value and isolated after precipitation as elemental selenium suitable for production of new filter mass.
3. A process according to claim 1 or 2, **characterised** in that the treatment with aqua regia is carried out at an elevated temperature.
4. A process according to any one of claims 1 to 3, **characterised** in that the mass which is freed of the aqua regia solution and only contains inert carrier material is transferred to production of new selenium filter mass after it has been washed and dried.
5. A process according to any one of claims 1 to 4, **characterised** in that the selenious acid formed during the hydrogen peroxide leaching is transferred to production of new selenium filter mass.

Fig. 1



INTERNATIONAL SEARCH REPORT

International application No.

PCT/FI 2004/000768

A. CLASSIFICATION OF SUBJECT MATTER

IPC7: B01D 53/64, C01B 19/00

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC7: B01D, C01B

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

SE,DK,FI,NO classes as above

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

EPO-INTERNAL, WPI DATA, PAJ

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	US 3786619 A (MELKERSSON ET AL), 22 January 1974 (22.01.1974) --	1-5
A	EP 0629431 A1 (BOLIDEN CONTECH AB), 21 December 1994 (21.12.1994) --	1-5
A	EP 0655794 A1 (BOLIDEN MINERAL AB), 31 May 1995 (31.05.1995) -- -----	1-5

☐ Further documents are listed in the continuation of Box C.

☒ See patent family annex.

* Special categories of cited documents:

- "A" document defining the general state of the art which is not considered to be of particular relevance
- "E" earlier application or patent but published on or after the international filing date
- "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- "O" document referring to an oral disclosure, use, exhibition or other means
- "P" document published prior to the international filing date but later than the priority date claimed

"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

"X" document of particular relevance: the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

"Y" document of particular relevance: the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

"&" document member of the same patent family

Date of the actual completion of the international search

20 April 2005

Date of mailing of the international search report

25 -04- 2005

Name and mailing address of the ISA/
Swedish Patent Office
Box 5055, S-102 42 STOCKHOLM
Facsimile No. +46 8 666 02 86

Authorized officer

Bertil Dahl/ELY
Telephone No. +46 8 782 25 00

INTERNATIONAL SEARCH REPORT

Information on patent family members

01/04/2005

International application No.

PCT/FI 2004/000768

US	3786619	A	22/01/1974	AU	463684	B	11/07/1975
				AU	4306772	A	13/12/1973
				CA	982497	A	27/01/1976
				DE	2227176	A	14/12/1972
				FR	2140238	A,B	12/01/1973
				GB	1389109	A	03/04/1975
				BE	784334	A	02/10/1972
				DD	96636	A	05/04/1973
				DK	130279	B,C	03/02/1975
				ES	403489	A	01/05/1975
				FI	51770	B,C	31/12/1976
				IT	956140	B	10/10/1973
				JP	55039364	B	11/10/1980
				NL	177189	C	16/08/1985
				NL	7207454	A	06/12/1972
				SE	360988	B	15/10/1973
				YU	34961	B	30/06/1980
				YU	147772	A	31/12/1979

EP	0629431	A1	21/12/1994	SE	502524	C	06/11/1995
				SE	9302111	A	19/12/1994

EP	0655794	A1	31/05/1995	SE	0655794	T3	
				AT	153480	T	15/06/1997
				AU	675587	B	06/02/1997
				AU	7742394	A	01/06/1995
				CA	2134731	A,C	26/05/1995
				DE	69403310	D,T	28/08/1997
				DK	655794	T	08/12/1997
				ES	2101479	T	01/07/1997
				FI	106131	B	00/00/0000
				FI	945560	A	26/05/1995
				JP	7185506	A	25/07/1995
				NO	301342	B	13/10/1997
				NO	944486	A	26/05/1995
				SE	9303905	A	26/05/1995
				US	5567223	A	22/10/1996